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Perspective

Nanomaterials Derived from Fungal Sources—Is It the New Hype?

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Supporting Information

MACROMOLECULES

ABSTRACT: Greener alternatives to synthetic polymers are constantly being investigated and sought after. Chitin is a natural polysaccharide that gives structural support to crustacean shells, insect exoskeletons, and fungal cell walls. Like cellulose, chitin resides in nanosized structural elements that can be isolated as nanofibers and nanocrystals by various top-down approaches, targeted at disintegrating the native construct. Chitin has, however, been largely overshadowed by cellulose when discussing the materials aspects of the nanosized components. This Perspective presents a thorough



overview of chitin-related materials research with an analytical focus on nanocomposites and nanopapers. The red line running through the text emphasizes the use of fungal chitin that represents several advantages over the more popular crustacean sources, particularly in terms of nanofiber isolation from the native matrix. In addition, many β -glucans are preserved in chitin upon its isolation from the fungal matrix, enabling new horizons for various engineering solutions.

INTRODUCTION

Fibers and filaments of different kinds have been utilized by humans for various materials, commodities, and engineering solutions throughout history. Natural fibers were complemented by synthetic fibers during the course of the 20th century, and nanofibers provided a new set of possibilities at the turn of the century. With polymers, synthetic nanofibers are generally prepared from a solution via electro¹ or solution blow spinning.² Carbon-based building blocks, conversely, have initiated a new field of research with, for example, carbon nanotubes, carbon nanofibers, and graphene nanoribbons.³ With a research scene populated by bottom-up approaches to obtain these aforementioned synthetic, nanosized filaments, the early 21st century has seen the rise of polysaccharide-based nanofibers, isolated from native sources via top-down methods. Cellulose nanofibers isolated from plant fibers have received substantial attention,⁴⁻⁶ followed by chitin nanofibers, predominantly from crustacean sources.⁷ In addition, shorter rod-like particles termed nanocrystals have been isolated both from cellulose⁸ and chitin,⁹ and their usage in diverse materials has also been subject to scrutiny.^{5,10} The research on native nanofibers and nanocrystals is above all driven by sustainability measures: renewability, biodegradability, and, in general, the

possibility to substitute fossil-based materials with greener solutions. Much of the research is based on the high strength and large surface area of polysaccharide nanomaterials as they originally provide the structural support in their natural environment. An added value is provided by the relatively high aspect ratio of cellulose and chitin nanomaterials, rendering them particularly suitable as a reinforcing phase in composites.^{11,12} Another branch of research, equally ambitious, focuses on more explicit properties of nanofibers and nanocrystals, as dictated by evolution: chirality,¹³⁻¹⁷ amphiphilicity,^{18,19} and specific response to water,^{20–23} for example.

The activity involving chitin nanofibers has always been slightly overshadowed by the efforts concerning cellulose nanofibers: the preparation requires additional steps and the nanofiber networks (nanopapers) or composites do not quite reach the same strengths as those reported for cellulose. In this

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Special Issue: Future of Biomacromolecules at a Crossroads of Polymer Science and Biology



Figure 1. Historical timeline for milestones in (nano)cellulose and (nano)chitin research.

Perspective, we aim at probing the true potential of chitin nanofibers with respect to their cellulosic counterparts. The focus is particularly set on chitin of fungal origin-a source that possesses distinct benefits over the current hegemony of crustaceans as the most common chitin supply. First, unlike the crustaceans, the fungi do not contain minerals that require an acidic extraction step for removal, thereby also partially degrading the chitin in the process. In fact, the isolation procedure for chitin nanofibers can be very simple indeed, requiring just brief mechanical agitation in a kitchen blender after a mild alkaline treatment to remove proteins.²⁴ Second, the fungi generally include a substantial amount of β -glucans, which may be advantageous for the subsequent materials prepared from chitin nanofibers. For instance, the authors of this Perspective recently showed that chitin nanopapers from a fungal source have superior tensile properties compared with their crustacean equivalents.²⁴ The main reason behind the increased strength was ascribed precisely to the presence of β glucans in the sample, imparting a composite character to the nanopapers.

We start by putting the discoveries with chitin in a historical perspective with cellulose research. Subsequently, we discuss chitin nanofiber and nanocrystal preparation and the principal target materials from those nanomaterials, namely, nanopapers, composites, and foams. These applications are particularly noteworthy within the modern discourse on plastic waste and the mitigation of plastic production in the first place. The red line running through the Perspective lies with the benefits of fungal chitin. In other words, we attempt to overturn the entrenched orthodoxy that has established crustaceans as the unchallenged source material for chitin.

BRIEF HISTORY OF (NANO)CHITIN VS (NANO)CELLULOSE

Figure 1 depicts the milestones in the field of chitin and cellulose research, including the major discoveries with their nanosized units. Although Braconnot's isolation of chitin preceded Payen's isolation of cellulose by almost 30 years, research and industrial application of chitin has lagged behind that of cellulose. An upsurge of interest in chitin only started in the 1970s, where at least three factors helped to contribute: (1) publication of "Natural Chelating Polymer"²⁶ and "Chitin" by Muzzarelli, (2) organization of first International Conference on Chitin and Chitosan in 1977,²⁷ and (3) growth of aquaculture and shellfish consumption in Asia.²⁸ Nowadays, the momentum continues with dedicated societies like the Japanese Society for Chitin & Chitosan (established in 1989), European Chitin Society (established in 1996), and Indian

Chitin Society (established in 2010). More and more dedicated symposiums have been organized across the globe.

We will now take a brief walk through the parallel history of chitin and cellulose research: the pioneering works and the breakthroughs (Figure 1). The overview is understandably superficial, and an interested reader can obtain additional details elsewhere.^{28–30}

In terms of fundamental order, the native structure of crystalline cellulose was under debate for most of the 20th century, initiated by the early X-ray studies by Sponsler and Dore in the 1920s,³¹ advancing significantly when the native crystalline form was found to consist of two distinct polymorphs (cellulose I_{α} and I_{β}),³² and culminating in the renowned works by Nishiyama et al.^{33,34} in the early 21st century where the structure of the two native polymorphs, I_{α} and I_{β} were laid down in 1 Å resolution. With chitin, the structures of α - and β -chitin were published in the 1960s,³⁵ refined in the 1980s,³⁶ and finally published in 1 Å resolution in 2011.^{37,38} In conclusion, the unravelling of the crystalline structure of chitin has very closely followed that of cellulose.

Unlike starch, which is granular in nature, both cellulose and chitin are fibrous. Hence, they are prime candidates for sheets or films, or for the reinforcing phase in composite materials. A stronger paper can be made by fibrillating (disintegrating) the fibers into nanofibers. Smaller fibers provide higher surface area, which consequently leads to a higher contact area in a composite matrix or stronger sheets in a paper-like material. Bacterial cellulose (BC) is a special case among cellulosic substrates because the nanosized fibrils are extruded by bacteria directly into an isotropic structure with no hierarchical morphology.³⁹ Therefore, BC can be viewed as nature's own species of nanocellulose in contrast to plant-based cellulose where the hierarchical fiber structure must always be disintegrated in order to isolate the nanofibers from the matrix. Although a number of accounts from 1940s onward had shown that harsh mechanical treatment, particularly ultrasonication, can disintegrate plant fibers into cellulose nanofibers,⁴⁰ a major breakthrough in nanofibrillation was published in 1983 where Turbak et al.⁴¹ demonstrated largescale homogenization of pulp fibers into nanofibers for usage in materials technology. However, heavy energy consumption and instrumental demands kept cellulose nanofibers out of the spotlight until around 2006-2007 when a string of publications emerged by groups in Europe and Japan, advocating the use of various chemical and enzymatic pretreatments to significantly facilitate the mechanical fibrillation.⁴²⁻⁴⁵ It was soon noticed that chitin nanofibers



Figure 2. Structure of chitin on different length scales, starting from an exemplary crustacean source (photograph, top right corner), moving onto electron microscope images before and after demineralization, and showing simplified schematics of proteins enfolding the chitin nanofibers, which consist exclusively of chitin polymers. Chitin can be subsequently converted to chitosan with deacetylation.

(ChNFs) can also be isolated by similar means from a variety of sources.^{46–48}

Nanocrystals represent another type of polysaccharide-based nanomaterials where the semicrystalline nature of native cellulose or chitin is utilized by selectively hydrolyzing the noncrystalline domains by strong acids while leaving the crystallites intact. Cellulose nanocrystals (CNCs) were discovered in ca. 1950 by Rånby,⁴⁹ but the interest in CNCs surged when the Gray group at McGill University (Canada) found out in 1992 that CNCs spontaneously arrange into chiral nematic liquid crystals in colloidal suspensions.¹³ Chitin nanocrystals (ChNCs), in turn, were first introduced in 1959 and received more attention after the turn of the century, although their tendency to form chiral nematic liquid crystals was established soon after the similar discovery with CNCs.¹⁴

Because BC is readily available in nanosized form without any additional treatments, Yamanaka et al.⁵⁰ could demonstrate already in 1983 that BC sheets possess a remarkable tensile modulus (E) and tensile strength (σ) of 15–18 GPa and ~250 MPa, respectively. Following the isolation studies of plant-based nanofibers, the first study on strong sheets from wood-based (chemical pulp) nanocellulose (nanopaper) was carried out in 2008 by Berglund's group in Sweden (E = 13.2 GPa, $\sigma = 214$ MPa).⁵¹ Later, by orienting the fibers by a means of cold drawing,⁵² they managed to produce the highest value for cellulose nanopaper so far: E = 24.6 GPa, $\sigma = 428$ MPa. This is 13 times stiffer and 13 times stronger than standard printing paper.

Predating the papers on actual ChNF isolation, the first report on chitin nanofiber sheets appeared in 1992,⁵³ in which sheets made from crab chitin (α -chitin) were compared to sheets made from squid pen chitin (β -chitin). However, the nanofibers used in that study had a relatively large diameter (~100 nm in width). Research on nanopapers with smaller chitin nanofibers (10–20 nm in width) started more recently in 2010 by a Japanese team led by Ifuku.^{7,46,47,54} Their chitin sheets had decent mechanical properties (E = 2.5 GPa, σ = 42 MPa).⁵⁵ Recently, Berglund's group⁵⁶ and the present authors²⁴ managed independently to obtain substantial improvement in chitin nanopaper strength (E ~ 7 GPa, σ ~ 200 MPa) which are the highest reported value for chitin sheets in the literature so far.

Concerning cellulose nanocomposites, pioneering works in 1995 by a group of scientists at French research institute CERMAV have led the way.⁵⁷ At first, they used CNCs to reinforce a polymeric latex matrix. By adding 6% tunicin CNCs⁵⁷ or 30% wheat straw CNCs,^{58,59} they observed a 1000-fold improvement in storage modulus. Later, they used CNFs to reinforce a starch-based matrix.^{60,61} By adding 5% potato pulp nanofibers, they observed a significant improvement (at least 2 orders of magnitude) in the storage modulus compared with a neat starch film.

Inspired by the reinforcing effect provided by cellulose, pioneering work on nanocomposite reinforced by chitin followed. ChNCs isolated from squid pen,⁶² Riftia tube,⁶³ and crab shells were used for a number of composite structures.^{64–66} The first report on ChNFs as a reinforcement in composites came out much more recently. In 2011, Ifuku et al. impregnated ChNF film with 11 different types of thermoplastic (meth)acrylic resin to obtain transparent nanocomposite films—all of which show great improvement in mechanical properties over the neat resin.⁵⁵

All the aforementioned nanocomposite studies deal with water-based or thermoplastic resins. Thermoset matrices, such as epoxy or phenol-formaldehyde, are often favored when a high-performance material is required. The first study on using tunicate CNCs to reinforce a waterborne epoxy resin was published in 2000.⁶⁷ However, a major breakthrough was made 5 years later by Yano et al. in Japan.⁶⁸ They demonstrated that a strong nanocomposite with a bending modulus (E_b) of 19 GPa and a bending strength (σ_b) of 370 MPa can be made when wood-based (kraft pulp) CNFs were used as a reinforcement for a phenolic resin. An even higher modulus and strength ($E_b = 28$ GPa, $\sigma_b \sim 410$ MPa) was obtained when

Table 1. Sources and Properties of α -Chitin and β -Chitin

	α-chitin	β -chitin
sources	shells of crustaceans lobster ⁷⁹ crah ⁴⁶	squid pen ⁸⁸ sea tubeworms ⁸⁹ centric diatom
	shrimp ⁸⁰ krill ⁸¹	$Thalassiosira^{90} \rightarrow most crystalline$
	cuticle of insects ⁸² cell wall of fungi	
	mushroom ⁸³ yeast ⁸⁴ mold ⁸⁵	
	marine alga $Phaeocystis^{86} \rightarrow highly crystalline$	
	grasping spine of arrow worms $Sagitta^{87} \rightarrow most crystalline$	
molecular packing	orthorhombic ^{86,91} (a: 4.75 Å b: 18.89 Å c: 10.33 Å x: 90°)	monoclinic ^{37,90} ($a: 4.82 \stackrel{h}{=} h: 9.25 \stackrel{h}{=} c: 10.39 \stackrel{h}{=} x: 97.2^{\circ}$)
chain arrangement	antiparallel; analogous to mercerized or regenerated cellulose (cellu	alose II) parallel; analogous to native cellulose (cellulose I)
hydrogen bonding	has intersheet bonding has intrasheet, interchain, and intrachain bonding	no intersheet bonding has intrasheet, interchain, and intrachain bonding
polymorph stability	stable	metastable, converted into α -chitin by 25–30% NaOH ⁹² 8% HCl ⁹³
thermal stability lattice modulus (axial)	isotropic lateral expansion when heated ⁹⁴ theoretical (calculation): ~150 GPa ⁸²	more easily deacetylated ⁸⁸ anisotropic lateral expansion ⁹⁴ when heated not reported
solubility	stable, not soluble in most organic solvent	soluble in formic acid ⁹⁶ susceptible to swell in water ⁹⁷
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Figure 3. α - and β -chitin structures with c axis representing the fiber direction. Adapted with permission from ref 101. Copyright 2006 Elsevier.

BC nanofibers were used in the same resin.⁶⁹ This is the highest reported value in the cellulose nanocomposite field so far, and to put this into perspective, the measured strength is almost comparable to structural steel. It is worth noting that Yano and co-workers used a high cellulose content (>80%) and a very high compaction pressure (50–100 MPa) to prepare their nanocomposite. Research involving nanoscale chitin in a thermoset matrix is much rarer. The first study was published in 2013, when Shao et al.⁷⁰ impregnated 40%

ChNFs with an epoxy resin. No data on mechanical properties were presented. Later in 2016, Shibata et al.⁷¹ reinforced their waterborne epoxy resin with 3% ChNFs, but their nanocomposite performance (E = 2.4 GPa, σ = 50 MPa) was nowhere near the cellulose nanocomposites prepared by Yano. The authors claimed that their nanocomposite performance was actually reduced at higher loadings of chitin. As for ChNCs in a thermoset matrix, we are yet to find a single study about it.

Table 2. Chitin Content in Selected Fungi Species^a

	per dry weight		per AIM ^b		
fungi	chitin (%)	ref	chitin (%)	glucan (%)	ref
A. bisporus (common mushroom)					
whole	3-9	83,107,108	36	18-36	110
stalk	7-19	107,109	34-44		109,111
cap	6-7	107		27	
mycelium			31		111
L. edodes (shiitake mushroom)	1-10	83,107,108	28	68	112
P. ostreatus (oyster mushroom)	2-15	107,108,113			
P. eryngii (king trumpet mushroom)	3-9	83,108,113			
S.commune (split gill mushroom)			22	68	114
A. niger (black mold)	8-27	115,116	24	40	115
S. cerevisae (baker's yeast)	1-3	117	3	37	118
M. rouxii (white mucor)	8-9	115,119			

^{*a*}The values from literatures were rounded to nearest decimal. Only the lowest and the highest value are presented. ${}^{b}AIM = alkali$ insoluble material.

Our brief walk into the history of cellulose and chitin shows one undeniable fact: progress in chitin follows the footsteps of cellulose. We can attribute this to the rich history of cellulose itself, spanning over millennia as a source for clothing, building materials, and energy. Abundance and readily available sources enabled the prosperity of textile and paper industry, which further catalyzed research in cellulose. When the numbers of scientific publication of cellulose, chitin, and chitosan were combined, Scopus database reveals the following order: cellulose (71%) > chitosan (19%) > chitin (10%). If chitin researchers can equip themselves with some knowledge about cellulose, at least by following the trends, they will be much better prepared in their own studies. In the coming sections, we will focus on chitin. Readers who are interested in nanocellulose and their nanocomposites are referred to a thorough monograph written by Dufresne⁷² and to a number of recent reviews.^{11,12}

STRUCTURE AND SOURCE OF CHITIN

Chitin is structurally similar to cellulose except that the C2hydroxl group of cellulose is replaced by an acetamide group (Figure 2). If this group is deacetylated, the polymer becomes chitosan. Primary amine groups in chitosan can be protonated to cations in dilute acid; thus, it is more soluble than chitin and can (1) confer antibacterial properties when used in hydrogel form,^{73,74} (2) chelate heavy metal ions and dyes in wastewaters more efficiently than chitin,^{75–77} and (3) be manipulated more easily than chitin for added functionality.⁷⁸ Given these factors, it is not surprising that more research has been performed on chitosan than on chitin. Nevertheless, the susceptibility of chitosan in dilute acid makes it unsuitable for applications that require durability such as strong films or composites.

Chitin exists in two major polymorphic forms, α and β . Their sources and related properties are summarized in Table 1, while their molecular packing is depicted in Figure 3. It has been suggested that the third polymorph, γ -chitin, may be a distorted version of either α - or β -chitin rather than a true third polymorphic form.⁹⁸ The similarity to the properties of cellulose is apparent from Table 1. The experimental modulus (ca. 60 GPa) of α -chitin falls short on the measured values for native cellulose I (ca. 110–220 GPa),⁹⁹ but the order of magnitude is the same. Compared with cellulose, chitin also has lower expansion coefficient and a higher degradation temperature.¹⁰⁰ Zooplankton cuticles (in particular Antarctic krill, with the estimated biomass of 379 million tons),¹⁰² constitutes the largest source of chitin on earth.¹⁰³ Waste from the shellfish industry (shrimp, crab, and lobster shells) in which the chitin content ranges between 8 to $40\%^{64,104,105}$ constitutes the main source of commercial chitin production nowadays. The chitin content in fungi is usually lower than in crustacean sources; the chitin contents for few selected species are listed in Table 2. In addition, Table 2 shows chitin and glucan contents for corresponding alkali extracted substrates as this is the regular step to remove proteins from the fungal matrix (see next section).

The primary biological function of crustacean chitin and fungal chitin is to provide structural support of the animal exoskeleton or fungal cell wall. However, this function is fulfilled differently because of their diverse physiochemical environments. Shellfish chitin normally binds with sclerotized proteins and minerals, while fungal chitin is associated with other polysaccharides such as glucan and mannan.¹⁰⁶

During the extraction process—even if the treatment is very harsh-not all nonchitinous content can be eliminated, suggesting the existence of covalent-like cross-links between chitin polymers and other substances. This is quite different than interaction between cellulose-hemicellulose-lignin in woody biomass. Cellulose is physically bonded (rather than covalently bonded) to the rest, while hemicellulose and lignin are purportedly covalently linked to each other.¹²⁰ Because of this, it is easier to obtain higher-purity cellulose than highpurity chitin, although even with cellulose it is virtually impossible to fully extract all noncellulosic materials from the plant fiber matrix. (BC is an exception here because it is biosynthesized solely as pure cellulose without additional ingredients.) In animal chitin, unless if the sources are from diatom Cyclotella or Thalassiosisira-which is pure chitin, no associated protein¹²¹—residual protein will always coexist with chitin.

In crustacean shells and insect cuticles, the cross-links between chitin and protein are well-known. However, it is still debatable whether the bridging is partially covalent in nature^{35,122,123} or not.^{124,125} Given that the amount of residual protein is very low, the amount of covalent bonding is probably low, or the bonds may be cleaved during the isolation process. Therefore, chitin contents from animal sources cited in the literature are generally reliable.



Figure 4. (a) Difference between glucan structures in fungi and cellulose. (b) Right: where chitin resides in mushroom; left: TEM image represents hyphal tip of *S. rolfsii*. Reprinted with permission from ref 139. Copyright 1988 Springer Nature. https://link.springer.com/journal/709.

In fungi, covalent linkages between chitin and glucan have been demonstrated by chemical hydrolysis and enzymatic dissection,^{126,127} gene disruption,¹²⁸ and solid-state NMR.¹²⁹ The initial study was done on *S. commune* (split gill mushroom),¹²⁶ but similar results were also found for other fungi.^{130–132} Unlike animal chitin where the residual protein is minimal, fungal chitin contains a higher proportion of glucan, often higher than that of chitin itself. Thus, any literature citing the percentage of chitin from fungi should be treated with caution, especially if the work is not directly related to mycology. More often than not, the stated chitin value represents the alkali insoluble material (AIM), in which chitin and glucan coexist. Alkali insoluble glucan exists in β -form, often called β -glucan.

There are numerous entries discussing fungal β -glucan and its structural diversity.¹³³ However, most of them refer to the water-soluble part only. For example, lentinan is β -glucan obtained from the hot water extract of *L. edodes* (shiitake mushroom), schizophyllan from *S. commune* (split gill mushroom), zymosan from *S. cereviase* (baker's yeast), pleuran from *P. ostreatous* (osyter mushroom), grifolan from *G. frondosa* (maitake mushroom), and ganoderan from *G. lucidum* (lingzhi or reishii mushroom). All these glucans were studied mainly because the human innate immune system can recognize them;¹³⁴ hence, they boast a remarkable potential in applications such as immune stimulator, antibacterial, antitumor, anticancer, antioxidant, and other health-related benefits. Because this Perspective is focused on the materials potential of chitin after purification, we were only interested in alkali insoluble β -glucan; thus, we will not further discuss the soluble glucans, and an interested reader is referred to critical reviews on the topic.^{135–137}

Structural intricacies and association with chitin makes β glucan in fungi partially insoluble. When the alkali resistant part of the cell wall is extensively treated with glucanase (glucan degrading enzyme), 16% of glucan still remained in the insoluble form.¹³⁸ However, when the same resistant part is treated with chitinase (chitin-degrading enzyme), all glucan became soluble.¹²⁶ Both of these results indicated a cross-link between chitin and glucan. Insolubility during the glucanase treatment is due to crystalline chitin, while total solubilization after chitinase treatment reflects the collapse of the crystalline structure. Thus, logic dictates, if some glucan is still in the insoluble form after extensive glucanase treatment, then it must somehow be linked to chitin. It is worth noting that the glucanase action is highly specific and was carried out on the alkali resistant part, so typical alkali extraction will leave more glucan connected to chitin.



Figure 5. Architecture of chitin-glucan complex in fungi. The mushroom model is based on *S. commune*,¹⁴⁰ the yeast model is based on *S. cerevisae* and *C. albicans*,^{84,127,141} and the filamentous fungi model is based on *A. fumigatus*.¹³¹

36

Insoluble β -glucan usually consists of $(1\rightarrow 3)$ backbone with $(1\rightarrow 6)$ branches, as opposed to the $(1\rightarrow 4)$ glycosidic linkages in cellulose (see Figure 4a) and hemicellulose. Its proportion and branching is highly dependent on the species and the extraction process. Harsh acid treatments will degrade most of the glucan, causing the X-ray diffraction pattern of the sample to resemble that of crustacean chitin.¹³⁸

The relation between chitin and insoluble glucan in the fungal cell wall is depicted in Figure 4b. Extensive studies have been carried out to elucidate the architecture of chitin-glucan linkages in Ascomycete phyla (yeast and filamentous fungi). However, similar studies with basidomycete (mushroom) are still lacking. Figure 5 illustrates the chitin-glucan model gathered from literature. There are small differences between the insoluble glucans among mushrooms, yeast, and filamentous fungi. Nevertheless, for the most part, all of them bear the common motif: β -glucan, which is associated with chitin, is predominately having a $(1\rightarrow 3)$ backbone with a $(1\rightarrow 6)$ branching.¹⁴² In yeast, most of the chitin is concentrated at the bud scar, while for the other fungi, chitin is present throughout the cell wall. All fungi synthesize chitin in their cell wall, but only zygomycete (mucor species) are known to be able to cosynthesize chitin and chitosan simultaneously.^{143,144}

ON THE USE OF FUNGI AS A POTENTIAL SOURCE FOR CHITIN

Recent estimates suggest that as many as 5.1 million fungal species might exist, but only fewer than 100 000 have been described so far.¹⁴⁵ Fungi grow by hyphal branching, creating a vast three-dimensional network of web-like mycelia. An individual hyphae is about $2-3 \mu m$ in diameter. Mycelium in itself has many prospects to be used as an alternative biorestoration medium, for example, as water filtration media for pathogens¹⁴⁶ or for bioremediation.¹⁴⁷ Certain species assemble their hyphae into more complex structures such as hyphal strands of many linearly arranged hyphae and particularly into large fruiting bodies (mushrooms and toad stools) in which the hyphae are arranged in various complex

formats and, typically, single or multicellular spores are generated for dispersal.

At the cellular level, different fungal species have different ratios of chitin to glucan in their cell walls. On one hand, the covalent linkage of amorphous glucan with chitin will complicate materials characterization, but on the other hand, it offers material scientists with new opportunities to explore and utilize a different class of renewable polymeric (natural) composite materials down to a nanoscale. While chitin is a rigid linear polymer, branched glucans act as amorphous matrix, which after extraction from mycelium or fruiting bodies results in a natural nanocomposite architecture with varying proportions of chitin and glucan optimized by nature. This is a feature of the chitin from the fungal species that does not exist for animal chitin.

Table 3 summarizes the pros and cons of chitin extracted from fungi and animal sources. Fungal chitin is free from the crustacean allergenic protein, tropomyosin,¹⁴⁸ which further extends its potential usability. For example, KitoZyme (www.kitozyme.com) has extracted a chitin-glucan complex from *Aspergillus niger* for marketing as a food supplement.¹⁴⁹ The scientific committee of the European Food Safety has regarded their chitin-glucan supplement as safe to use,¹⁵⁰ and the U.S. Federal Drug Agency also approved their GRAS (Generally Recognized as Safe) status. In the U.K., Quorn products (www.quorn.co.uk) utilize mycoprotein extracted from *Fusa-rium venetatum* as a popular meat substitute.

However, harnessing fungal chitin is not without some challenges. The chitin yield per wet weight of mycelium or fruit body is relatively low compared with the animal-based counterpart. This can be mitigated to a large degree by the rapid growth rate of many fungi on a diversity of simple substrates or "waste" organic matter and relatively simple extraction protocols. However, the major *perceived* obstacle discouraging researchers from working with fungal chitin has been the presence of glucans. In contrast with this conventional wisdom, we contend that, in fact, this represents a huge opportunity with considerable innovation and exploitation potential.

Table 3. Pros and Cons of Crustacean-Based and Fungal-Based Chitin^a

chitin source	advantages and disadvantages			
crustacean	high chitin content per dry mass			
	• almost pure chitin \rightarrow easy to characterize			
	• values put e child \rightarrow easy to characterize • wall established extraction and nanofiber preparation protocols			
	renewable resource			
	 already commercialized and can be bought easily 			
	extensive research and literature accounts			
	Disadvantages			
	• limited supply \rightarrow seasonal and regional			
	• sea pollution can affect chitin quality			
	• problems associated with the crustacean allergenic protein "tropomyosin"			
	• demineralization step requires the use of concentrated acid			
fungi	Advantages			
-	• renewable, nonallergenic			
	• not dependent on seasonal fluctuation			
	• fungi			
	-do not need sunlight; can be grown anywhere, anytime			
	-fast growth (mushroom) \rightarrow typically 2–3 weeks			
	-can be grown by stacking (vertical growth) \rightarrow minimal land required			
	-year round supply			
	-growth can be controlled easily \rightarrow more consistent chitin quality			
	-inexpensive raw material for growth, organic waste can be used as a growth substrate			
	-zygomycetes \rightarrow direct chitosan extraction, no deacetylation process required			
	• readily available fungal biomass, e.g., "A. niger" from industrial citric acid production			
	• readily available mushroom waste (irregular shape and stalk) from mushroom industry			
	• no demineralization step required			
	- reduces overall production cost			
	- no chain depolymerization due to acid treatment			
	• night level of acetylation on childh \rightarrow offers a certain level of hydrophobicity • may proceed bioactive function, due to presence of β always			
	• may possess bloactive function, due to presence of p-glucan			
	Disadvantages			
	low chitin content per wet weight			
	• not pure chitin, association with glucan \rightarrow complicates characterization			
	• extraction process not vet widely demonstrated at industrial level (with a few exceptions)			
	Imited literature on extraction for nanofiber production			
	t			

 a Input were gathered from ref 151, other literature, and our own observations.

In comparison to nanocellulose, the chitin in fungi requires also a special notice. The cellulose content in trees, for example, is known to be ca. 40–50%, which is well above the dry weight content of chitin in fungi (Table 2). However, it takes years or even decades for a tree to grow to such state that it can be harvested for nanocellulose, whereas it is just a matter of weeks before chitin nanofibers (or nanocrystals) can be extracted from the native source. Besides, the isolation of nanochitin from fungi is far more effortless than the isolation of nanocellulose from wood (see the sections below). Another advantage is the inclusion of β -glucans with the nanochitin structure, eventually enabling stronger nanopaper construction among other benefits.

Isolation of Chitin and Nanosized Chitin. In nature, chitin coexists in a matrix with other materials that are usually minerals, proteins, and glucans, depending on the native source as described in the previous section. In order to isolate the actual ChNFs or ChNCs, the source material must generally consist of more or less pure chitin, and the use of several extraction steps is required. Here, the fungal chitin is again in a more favorable position: the extraction is simpler and the

isolation of ChNFs in particular is more effortless with minimal energy consumption.

Extraction of Chitin from the Native Source. There are three main steps usually required to extract chitin from crustacean shells: demineralization, deproteination, and decoloration.¹⁵² A major advantage with fungi is that they are not calcified; thus, the demineralization step is unnecessary, although deproteination must be carried out.^{153–155} A deacetylation step is subsequently added if a chitosan product is preferred (Figure 6).

Demineralization removes inorganic material in calcified crustacean shells. It is usually carried out using dilute acid, commonly hydrochloric acid (HCl), at room temperature. Care must be taken as acid can both deacetylate and depolymerize the chitin chain.^{156,157} In his optimization study, Percot et al.¹⁵⁸ reported that shrimp shells are completely demineralized within 15 min of 0.25 M HCl treatment at room temperature. Longer treatment times, higher concentrations of acid, or higher extraction temperatures will all cause a substantial reduction in chitin molecular weight. Although HCl is relatively expensive and detrimental to chitin, it remains the most commonly used decalcifying



hexafluoro-2-propanol, CaCl2/MeOH = calcium chloride/methanol, TCA/DCE = trichloroacetic acid / 1,2-dichloroethane, NaCH+CS₂ = sodium hydroxide + carbon disulfide (xanthate or viscose process), TEMPO = 2,2,6,6-tetramethylpiperidine-1-oxyl

Figure 6. Flowchart for isolation of chitin and chitosan from crustacean shells and fungi, and preparation of ChNF and ChNC.

agent in both laboratory and industrial-scale production of chitin. Weak synthetic amino acids like ethylenediaminetetraacetic acid (EDTA) at controlled pH have been proposed as nondegradative demineralization reagents.¹⁵⁹ However, elimination of inorganic salts was found to be incomplete.¹⁵²

Deproteination, that is, the removal of protein, is a crucial step in chitin extraction for both crustacean and fungal sources. Sodium hydroxide (NaOH) and potassium hydroxide (KOH) are preferred reagents. They are typically used at 1 M concentration with variations in temperature and extraction time. Deproteination is less damaging to chitin compared with demineralization, but prolonged treatments and high temperatures can lead to deacetylation.¹⁵⁸ Higher deacetylation leads to (1) decrease in hydrophobicity, (2) decrease in nanopaper or film tensile strength, (3) increase in film ductility, (4) increase in solubility, (5) increase in cell adhesion and proliferation, and (5) increase in the rate of biological degradation.^{160–163} The more deacetylated the chitin, the more it resembles chitosan. The use of proteolytic enzymes¹⁶⁴ provides an alternative to the harsh chemical treatments, minimizing the effect of deacetylation and depolymerization. However, there are cost-related problems associated with enzyme usage. To bring down the production costs associated with the deproteination step, Ifuku et al.¹⁶⁵ forewent the whole process and came up with a chitin-protein nanofiber product. They found that composite films (with acrylic) have almost similar mechanical properties compared to the composite made from fully deproteinated chitin nanofibers.

Greener and more cost-effective extraction processes can be achieved via biological fermentation.¹⁶⁶ For example, Jung et al.¹⁶⁷ used lactic acid producing bacteria, *Lactobacillus* *paracasei*, to produce acid for the demineralization step, and protease producing bacteria, *Serratia marcesens* to remove protein during their deproteination step. Compared with the chemical process, biological treatment can result in higher crystallinity and higher molecular weight of the chitin or chitosan product.¹⁶⁸ Recently, Boric et al.¹⁶⁹ introduced a completely different, solvent-free approach for deproteinization for crustacean waste, based on dielectric barrier discharge plasma. The method appears greener than the solvent-based techniques, but there is still room for optimization as a substantial part of proteins still remained within the chitin matrix.

Acid and alkaline treatment alone produces colored chitin. Hence, a decoloration step is added when a bleached product is desired. Pigment can be removed using ethanol or acetone after the demineralized step. Alternatively, the chromophores may be removed by bleaching using sodium hypochlorite (NaOCl) or an oxidation process using hydrogen peroxide (H_2O_2) .

The deacetylation step of chitin into chitosan is usually achieved by treating chitin with 40–50% NaOH at 95–100 °C for 2–3 h, followed by neutralization. The chitosan is then extracted with 2% acetic acid solution, filtered, and precipitated in distilled water. If the deacetylation process is carried out at room temperature, it yields a water-soluble form of chitin (i.e., alkali-chitin) instead of chitosan.^{160,170–172}

Isolation of Chitin Nanofibers and Nanocrystals. In recent years, a more intensive focus has been set on chitin nanomaterials, either as ChNFs⁷ or ChNCs.⁹ Just like native cellulose resides in nanosized "microfibrils" in plants, native chitin resides in nanosized fibers in the structural scaffold of

Table 4. Diameter of Chitin Nanofiber from Different Nanofibrillation Process

source	method	width (nm)	ref
crab shell	grinding + homogenization	<50	187
	grinding, pH 3–4	10-20	47
	microfluidizer	20-30	188
	ultrasonication, 24 kHz, 120 min, pH 3–4	2-20	189
	20% NaOH \rightarrow grinding, pH 3-4 ^{<i>a</i>}	10	190,191
	water jet atomizer "Star Burst"		
	(5 or 10 pass), neutral pH	17.3–18.2	181
	(1, 5, or 10 pass), pH 3	16.5-19.0	181
	high-speed blender, 10 min		
	37000 rpm, neutral pH	77 ± 37	192
	37000 rpm, pH 3-4	20-30	192
	15000 rpm, pH 3-4	20-30	193
	11000 rpm, pH 3-4	20-30	193
	4000 rpm, pH 3-4	~100	193
shrimp shell	grinding, neutral pH	10-20	80
	ultrasonication, 60 kHz, 30 min + pulse sonication	20	184
	electrospinning	670–µm	194
	domestic blender, 30 min, pH $3-4^{b}$	<50	195
squid pen	grinding, pH 3	12-20	196
	ultrasonication, 19.5 kHz, 2 min, pH 3–4	3-4	197
	self-assembly ^c		
	dissolution in HFIP \rightarrow solvent evaporation	3	198,199
	dissolution in LiCl/DMAc \rightarrow precipitation	10	198
lobster shell	homogenization	80-100	179
mushroom	grinding, pH 3	20-28	83

^{*a*} α -chitin is partially deacetylated with 20% NaOH (the surface of nanofiber behave like chitosan, but its core is chitin) followed by grinding in acidic condition. ^{*b*}Typical maximum rotational speed for conventional food mixer or kitchen blender is between 11000–15000 rpm. ^{*c*}HFIP = 1,1,1,3,3,3-hexafluoro-2-propanol, LiCl/DMAc = lithium chloride/N,N-dimethylacetamide.

crustaceans, insects, or fungi. These structural units can be isolated as nanofibers, resulting in ChNFs, or they can be partially degraded into shorter, rod-like ChNCs. Regardless of whether dealing with ChNFs or ChNCs, most studies utilize animal-based chitin as their starting material. To the best of our knowledge, only a few studies related to the deliberate extraction of fungal-based chitin nanomaterials (FChNFs or FChNCs) have been conducted so far.^{24,173–175} However, a number of older studies exist where mushroom pulp, prepared in a kitchen blender, has been used for papermaking.^{176–178} Because of the ease of ChNF isolation,^{24,173} these papers consist probably of chitin microfibers or even FChNFs—quite possibly with proteins and glucans—although their micro- or nanoscopic morphology was never reported.^{176,177}

Table 4 summarizes the width of ChNFs obtained from different nanofibrillation processes. All nanofibers in the table are several micrometers in length. ChNFs can be prepared by subjecting extracted chitin to various mechanical nanofibrillation processes such as high pressure homogenization,¹⁷⁹ wet shear grinding,¹⁸⁰ water jet atomization,¹⁸¹ high-pressure water jet with grinding pretreatment,¹⁸² microfluidization,¹⁸³ ultrasonication,^{184,185} or high-speed blending.¹⁸⁶ All these treatments are similar to nanofibrillation treatments used for cellulose. All nanofibrillation processes, except for ultrasonication, rely on high shearing and high impact force generated onto a chitin fiber bundle, causing weak interfaces among nanofibers to be broken. In ultrasonication, on the other hand, high frequency oscillation creates a localized highpressure region, resulting in cavitation and impaction, ultimately loosening the fibers.

Chitin nanofibers can also be prepared by a chemical method via TEMPO-mediated oxidation, similar to the conditions with cellulose, that is, in the presence of a catalyst (TEMPO), sodium bromide (NaBr), and an oxidizer, namely sodium hypochlorite (NaClO).^{200,201} NaClO with TEMPO oxidizes exclusively the primary C6-hydroxyl groups of polysaccharides into carboxylic acid moieties via an aldehyde intermediate. The charges brought in by the carboxylates promote anionic electrostatic repulsion that separate individual fibers apart. This method of nanofibrillation was first demonstrated for cellulose by Isogai et al. in 2006.⁴² Total oxidation of native cellulose cannot be achieved by TEMPO even after addition of a large amount of NaClO,²⁰² but in the case of chitin, total oxidation can occur;^{203,204} hence, the oxidation process for chitin must be strictly controlled. When utilizing TEMPO for chitin, the outcome varies widely depending on the chitin source. TEMPO for squid pen β chitin produces neither ChNFs nor ChNCs,⁴⁸ TEMPO for highly crystalline tube worm β -chitin produces ChNFs (20–50 nm in width, several microns in length),48 and TEMPO for crab α -chitin produces only ChNCs (8 nm width, 340 nm in length).²⁰⁵ There are parallels in such behavior to cellulosic substrates where, for example, TEMPO oxidation of microcrystalline cellulose or dissolving pulp coupled with heavy sonication leads to CNCs instead of CNFs.²⁰⁶ In the latter case, the few glycosidic bonds remaining in the disordered regions of dissolving pulp microfibrils are probably cleaved during the TEMPO-oxidation conditions, ending up in CNCs after the isolation with sonication. Recently, another oxidation method with ammonium persulfate was reported to yield ultrafine (2-4 nm width) ChNFs from squid pen chitin.²⁰⁷ It

source	method	length (nm)	width (nm)	ref
crab shell	3 M HCl, 1.5 h, boiling	100-600	4-40	64
	3 M HCl, 6 h, boiling	100-500	10-50	214
	3 M HCl, 3 h, boiling	300	20	215
	33% NaOH \rightarrow pH 3-4 ^{<i>a</i>}	250 ± 140	6.2 ± 1.1	211
	64% H ₂ SO ₄ , 60 °C, 1.5 h	426 ± 10	$\sim 15^{e}$	216
	TEMPO oxidation	676 ± 13	$\sim 15^{e}$	216
	ammonium persulfate, 75 °C, 16 h	486 ± 52	~15 ^e	216
	ammonium persulfate, 75 °C, 16 h	400-500 ^d	15	217
	TEMPO oxidation	50-500	8-10	218
	TEMPO oxidation	250	15	219
	TEMPO oxidation	150-500	20-55	220
	O ₂ /laccase/TEMPO	480 ± 200	24 ± 17	221
	ionic liquid/methanol	several 100	20-60	222
shrimp shell	3 M HCl, 1.5 h, boiling	150-800	5-70	223
	3 M HCl, 6 h, reflux at 120 $^\circ \text{C}$	231-969	12-65	224
	3 M HCl, 1.5 h, 90 °C	200-500	10-15	225
	3 M HCl, 1.5 h, 90 °C	160 ± 77	16 ± 5	226
	3 M HCl, 1.5 h, 90 $^\circ \mathrm{C} \times$ 3	230-260	9-10	227
	choline chloride-ZnCl ₂ /acetic acid	100-700	30-80	228
squid pen	3 M HCl, 1.5 h, boiling	50-300	10	62
<i>Riftia</i> tube	3 M HCl, 1.5 h, boiling	500-10000	18	229
commercial powder	choline chloride-PTS ^b	200-400	12-44	230
	TEMPO /NaClO /NaClO	200 600	6-15	231
	TEMPO/NaCiO/NaCiO ₂	200-000	0-13	201

Table 5. Dimensions of Chitin Whiskers Isolated by Different Processing Methods

^{*a*} α -chitin is partially deacetylated by 33% NaOH (i.e., the crystallite surface behave like chitosan, but its core still chitin) followed by 1 min ultrasonication at pH 3–4 to promote cationization. ^{*b*}*p*-Toluene sulfonic acid. ^{*c*}Consists of β -chitin. ^{*d*}Bimodal distribution. ^{*c*}Width not explicitly defined.

is also possible to increase the surface charge of chitin by esterification with maleic anhydride to facilitate ChNF isolation.^{208,209}

In 2008, Fan et al.²¹⁰ reported a much simpler and faster way to produce ChNFs from squid pen β -chitin (3-4 nm in width and several micron in length) by simply sonicating a specimen for 2 min under acidic conditions. The idea was based on cationization of free amine groups on the chitin crystallite surface at pH 3-4. Cationization leads to electrostatic repulsion, similar to what happens during TEMPO oxidation; but in this case, it is a cationic repulsion instead of an anionic repulsion. In the aforementioned study, sonication works with squid pen β -chitin, but it did not work with tubeworm β -chitin or crab α -chitin, arguably because of higher crystallinity. Crab α -chitin can be fibrillated into uniform nanofibers with 10-20 nm diameter by subjecting a neverdried sample to a grinding treatment.¹⁸⁰ Drying causes the fibers to collapse and they lose their swelling capability, thus making it harder to defibrillate. Later it was found that it was possible to obtain nanofibers of similar width using dried chitin by a means of grinding in acidic conditions.⁴⁷ Treatment with 33% NaOH causes α -chitin to be partially deacetylated, resulting in chitosan-like surface with a chitin core. Because chitosan is protonated in aqueous acid, nanofibers from partially deacetylated chitin can be easily individualized at pH 3-4 as a result of cationic repulsion of the positively charged amino group.²¹¹ An alternative approach to controlled deacetylation was introduced recently by Ye et al.,²¹² namely the use of chitin deacetylase enzyme that enabled effortless isolation of ChNFs with widths between 25 and 45 nm. The degree of deacetylation could be controlled with the addition of different metal ions.

In 2010, the Rolandi group¹⁹⁸ found that when squid pen β chitin was dissolved in HFIP, it can reassemble itself into ChNFs (α -chitin) (3 nm width) during solvent evaporation. β chitin dissolved in LiCl/DMAc can also self-assemble during precipitation process, but it produces larger diameter nanofibers (10 nm width). With another bottom-up approach, Zhu et al.²¹³ managed to prepare chitin nanofibers of 27 nm width by using NaOH/urea solvent with a phytic acid coagulation bath.

ChNCs (also known as chitin nanowhiskers or chitin crystallites) are usually prepared by boiling a chitin sample in hydrochloric acid (HCl) followed by sonication. ChNCs form stable colloidal suspensions because of the presence of cationic surface charge on its crystallite surface, and they are able to form chiral nematic liquid crystals after reaching a critical concentration.¹⁴ With cellulose fibers as a source, stable colloidal CNCs require the use of sulfuric acid (H_2SO_4) instead of HCl because anionically charged sulfate groups are introduced as half-esters on the CNC surface during the hydrolysis.²³³ Although the presence of sulfate groups on cellulose crystallites induce anionic electrostatic repulsion, they are also detrimental for the thermal stability of cellulose.²³ This is one of the advantages of chitin over cellulose in the nanocrystal form, as the use of HCl does not affect the thermal stability of chitin. The yield and the dimensions of either ChNCs or CNCs is highly dependent on the acid concentration and the duration of hydrolysis. Higher acid concentrations and longer treatment times will cause substantial reduction in nanocrystal length-to-width ratio.²³⁵

TEMPO-mediated oxidation is another method for ChNC production. This method offers several advantages over the conventional acid hydrolysis: (1) the process is more



Figure 7. Tensile properties of chitin nanocomposites as a function of chitin weight fraction reported in the literature. As a comparison, the tensile properties of polyethylene terephthalate (PET), polypropylene (PP) and high-density polyethylene (HDPE) are included into the same figure. These values are obtained from Matweb (http://www.matweb.com). All references for the data points are mentioned in the text, referring to the figure.

controllable by the amount of oxidizer added, (2) ChNC recovery can reach 90%, and (3) no deacetylation of chitin occurs during TEMPO-mediated oxidation.9 Furthermore, partial deacetylation with TEMPO-oxidation was shown to result in amphoteric ChNCs.^{219,231} Ammonium persulfate is another oxidant that can be used for ChNC isolation,²¹⁷ usually ending up with a higher charge than with TEMPObased ChNCs.²¹⁶ Surveying a completely different bottom-up approach, Kadokawa et al.²²² found that chitin regenerated from an ionic liquid can reassemble into a whisker form when the resulting chitin-ionic liquid gel is soaked in methanol. Another example of using a different solvent to obtain ChNCs was shown recently by Hong et al.²²⁸ who managed to prepare ChNCs from shrimp shell chitin by using a deep eutectic solvent (DES), specifically choline chloride-ZnCl₂. By contrast to the route via ionic liquids, however, the DES mixture did not dissolve chitin but it was used as a hydrolytic medium with acetic acid. Another recent account describes the use of DES for ChNCs, cholin chloride-*p*-toluene sulfonic acid where the latter is used as a catalyst for hydrolysis.²³⁰

Interestingly, when β -chitin was used as a source for ChNCs, the resulting particles were spherical instead of rod-like.²³² Despite the entirely different shape, however, the β -chitin was transformed to α -chitin on the course of harsh acid hydrolysis to acquire ChNCs. Table 5 summarizes the dimensions of ChNCs obtained from different processing methods.

The following sections constitute the core of this Perspective, namely, an assessment on how ChNCs and ChNFs fare in their use in composites, nanopapers, and foams, and why they are important in the first place.

APPLICATIONS OF NANOSIZED CHITIN

From the seas to the skies, the Earth is slowly being choked by the effects of the so-called "plastic epoch", which has generated 8.3 billion metric tonnes of virgin polymer to date.²³⁶ Annual global polymer production has experienced prolific and continuous growth from 1950, when 1.5 million tonnes of polymers were produced, to volumes of 350 million tonnes produced in 2017.^{237,238} This rise in polymer production combined with a global shift from reusable to single-use products has seen packaging become the largest contributor to plastic waste, which now plasters every corner of the globe. Approximately 42% of all nonfiber plastics were used for packaging in 2015, with 54% of those plastics disposed of in the same year.²³⁶ Devastatingly, almost 80% of all plastic waste ends up in landfill or dispersed in the environment, with 12 billion metric tonnes of plastic waste predicted to be present in landfill or the environment by 2050.²³⁶ With plastic taking up to 1000 years to degrade, one questions the logic of producing



Figure 8. Tensile modulus and strength of chitin nanocomposites as a function of strain-at-failure. The values for PET, PP, and HDPE are obtained from Matweb. All data points are referred to in the text.

single-use materials, used almost exclusively for protecting and transporting consumer products for short periods, from a material that is so difficult to safely return to the environment.

One short-sighted solution by developed countries to the growing problem with plastic waste has been to export the problem to second and third world countries. However, on December 31st, 2017 China introduced the policy known as "National Sword", which abruptly ended the import of postconsumer plastic waste. China will only accept postconsumer plastic waste with a high standard of cleanliness. As a result, less than 10% of the postconsumer plastic waste generated in the G7 countries was exported to China, compared with the first half of 2017 where approximately 60% of this waste was exported to China. Consequently, this plastic waste now accumulates in landfill or is incinerated, which places a significant burden on the environment. This sparked renewed interest into the development of biobased solutions to replace fossil-based polymeric materials.

As chitin is one of the most abundant biobased materials after cellulose and can be easily extracted from the waste shell of crustaceans or from fungal biomass, significant research effort has been poured recently into developing chitin-based materials for the replacement of fossil-derived polymers to curb the global issue of plastic waste, particularly single-use plastics.²³⁹ Genuine industrial efforts are also budding at the moment. For example, Shellworks group, a young start-up from London, is currently developing technologies to convert

chitin from lobster shells into plastic plant pots and plastic bags.²⁴⁰ CruzFoam, a company based in California, is developing sustainable surfboards out of chitin, as well as chitin foams to replace expanded polystyrene foams used for packaging.²⁴¹

Chitin as a Reinforcement for Polymer Nanocomposites. A common theme in the introduction of most research articles focusing on chitin nanocomposites is the need to develop sustainable materials for a greener future. As ChNFs possess high tensile properties and can be derived from fungi, crustacean waste or other bioresources, the resulting chitin nanocomposites could serve as a renewable alternative to many fossil-derived polymers. In this section, we will investigate precisely this potential of ChNFs or ChNCs composites to compete with contemporary synthetic commodity polymers. Chitin nanocomposites are a relatively large field of research with a more comprehensive treatise on literature featured in Table S1 (Supporting Information).

Figure 7 summarizes the tensile properties of chitin nanocomposites reinforced with either ChNFs or ChNCs reported in the literature. The polymer matrices reinforced with ChNFs in this figure include thermoplastics from soy protein,²⁴² polylactide^{243,244} and plasticized polylactide,^{245–247} polyurethane,²⁴⁸ citrate-based bioelastomer,²⁴⁹ maize starch,²⁵⁰ potato starch,²⁵¹ thermoplastic starch,²⁵² polycaprolactone,²⁵³ acrylic,^{165,254} and biobased epoxy resins.⁷¹ Significant variation in the tensile properties of chitin nanocomposites can be

observed, even for the same chitin loading fraction (Figure 7). This could be due to batch-to-batch variation of chitin, as chitin can be extracted from multiple sources using very different extraction conditions (see previous sections). Nevertheless, it is clear that chitin nanocomposites with tensile moduli and strengths as high as 4.5 GPa and 96 MPa, respectively, at a chitin loading fraction of 70 wt % can be manufactured.²⁵⁴

In Figure 7, we also overlay the tensile properties of PET, PP, and HDPE as our benchmark synthetic commodity polymers for comparison. We can see that a significant amount of reported tensile properties of chitin nanocomposites performed worse than our benchmark polymers. One should emphasize that these chitin nanocomposites are mainly using starch-based polymer matrices or softer polymer matrices, such as polycaprolactone or flexible polyurethane. On a more positive note, Figure 7 also shows that chitin can indeed be used as reinforcement to produce chitin nanocomposites with tensile properties similar to our benchmark commodity polymers. When the loading fraction of the chitin exceeds 50 wt %, the properties of the chitin nanocomposites did even exceed those of the benchmark polymers. The existence of such threshold loading fraction is also observed for nanocellulose-reinforced polymer composites,11 albeit lower, around 30 vol %, presumably due to higher tensile properties of single cellulose nanofibers.

In addition to tensile modulus and strength of a material, an important design consideration for polymeric materials is its fracture resistance or fracture toughness. To the best of our knowledge, the fracture resistance of chitin nanocomposites has not been well studied. To compare chitin nanocomposites to our benchmark polymers, we use the strain-at-failure as an approximation. The tensile moduli and strengths of various chitin nanocomposites plotted as a function of their reported strain-at-failure are summarized in Figure 8. In the same figure, we have also overlaid the properties of our benchmark PET, PP, and HDPE. It can be seen from this figure that very few chitin nanocomposites compare with the strain-at-failure (and by assumption also fracture resistance) of PP and HDPE but some chitin nanocomposites did compare with PET in terms of strain-at-failure but unfortunately not in terms of modulus or strength.

While the tensile modulus and strength of chitin nanocomposites could match that of PET, PP, and HDPE, their fracture resistance (estimated through the strain-at-failure) did not. This is postulated to be due to the brittle nature of chitin nanofibers, which when used as reinforcement for polymers, will produce brittle chitin nanocomposites. Nevertheless, some chitin nanocomposites are able to match the performance (in terms of tensile modulus, tensile strength vs strain-at-failure) of our benchmark polymers. These chitin nanocomposites were produced using plasticized polylactide as the polymer matrix. It should be noted, however, that the chitin loading fraction in these nanocomposites was only 1 wt %. Even though the loading fraction of chitin is low, it is sufficient to increase the tensile modulus of plasticized polylactide four-folds, with a minor decrease of the strain-at-failure from 305% to the observed 262%.

Our simple comparative analysis pointed to the discrepancy between expectation to produce chitin nanocomposites with attractive properties (as renewable options to replace single use "plastics") and the actual achievements. Significant effort is still needed to utilize chitin, a variable waste-derived material with high tensile properties, as efficient reinforcement for renewable, degradable to produce sustainable alternative for a greener future, with a particular focus not only on tensile modulus and strength but also the composites' fracture resistance and potentially dynamic mechanical properties, such as low energy impact strength.

Chitin Films, Sheets, and Nanopapers. While chitin might not serve as excellent reinforcement for polymers to produce truly green sustainable nanocomposites, an efficient way of utilizing chitin is to produce highly agglomerated chitin nanofiber networks, for example, chitin nanopapers, sheets, and films. (Note: we use "film" to describe a potentially pinhole free thin material with an appearance of a polymer film, while "sheet" is used to describe a thin CNC networks, whereas we reserve the term "nanopaper" for nanocellulose (CNF or BC) and ChNF networks.) Generally, chitin nanopapers, sheets or films can be produced using three different methods: (1) by solvent casting, which involves the dissolution of chitin in appropriate solvents followed by regeneration (with E up to 2 GPa and $\sigma = 93.1 \text{ MPa})^{255}$ or in analogy to papermaking using (2) a ChNF or (3) ChNC suspensions, which results in chitin nanopapers or sheets. The first method produces regenerated chitin films without any defined fiber morphology. However, pure chitin papers can also be produced using regenerated chitin fibers.²⁵⁶ Already in the 1970s, researchers started exploring (partially deacetylated) crab chitin as paper additive. 257 Their success stimulated Rice to explore using fibers from mushroom extract for the preparation of mushroom papers.^{176,177} Apparently, papermaking with mushroom fibers was first introduced in 1985 at the third International Fungi and Fiber Symposium. In the early years, researchers were interested in mushroom papers because of the chemicalfree nature of the pulping process and their colorful appearance.¹⁷⁸ Nanosized chitin is not exactly applied in these premature accounts, but the material likely consists of micron-sized chitin fibers with an uncharacterized amount of proteins and glucans originating from the raw material. Nevertheless, these studies serve as an early reminder that the fungal matrix is superior to the crustacean counterpart as a resource for papermaking.

Much more recently, chitin nanopapers and sheets have attracted some interest; they are typically prepared from ChNF or ChNC suspensions, which are dewatered either by simply letting the water evaporate following solution casting, or by filtration followed by hot pressing, or by using dedicated sheet making equipment (for instance Rapid-Köthen). Chitin nanopapers (from ChNFs) usually have better mechanical properties than sheets made from ChNCs as result of more extensive fiber entanglement (Figure 9). More recently, the authors of this Perspective revisited the preparation of mushroom papers.²⁴ It is indeed possible to extract highquality fungal nanofibers (FChNFs) from mushroom extract after a mild alkaline extraction and low-energy blending. In contrast to ChNFs extracted from crustaceans, which are often still associated with residual proteins, FChNFs are carrying significant amounts of covalently linked amorphous glucans. The benefit of the glucan-linked to the ChNFs is that in this case, FChNF nanocomposite films consisting of varying amounts of ChNFs reinforcing an amorphous glucan matrix can be readily prepared by simple filtration followed by hot pressing. The mechanical properties of the FChNF films depend on their chitin and glucan content, which depends on



Figure 9. Comparison between the tensile properties of bacterial (BC) and nanofibrillated cellulose (CNF) (data from¹), chitin (ChNF) nanopapers (data from^{188,55,191,24,259–261,56,262–264}), sheets of chitin nanocrystals (ChNC) (Data from^{265,222}), and fungal chitin nanofiber (FChNF) films (data from^{24,258}) as well as regenerated chitin films (data from^{255,266,267}).

the part of the fungi and species from which they were extracted and enclose a wide property window.^{24,258}

The tensile properties of ChNF and FChNF papers and sheets made from ChNCs, regenerated chitin films are summarized in Figure 9 as well as in Table S2 (Supporting Information). Also within the realm of nanopapers, nanocellulose-based building blocks have received much more attention than ChNFs and ChNCs in the past two decades.^{11,268} Cellulose nanopapers have outstanding mechanical properties, and thus, we chose to juxtapose the reported properties of FChNF films and ChNF nanopapers with the properties of CNF and BC nanopapers. When looking at the nanocellulose-based papers, by and large, BC nanopapers outperform CNF nanopapers in terms of strengths and moduli. ChNF nanopapers and FChNF films have typically significantly lower moduli as compared with cellulose nanopapers. The strengths of the FChNF films produced by mild alkaline extraction and low-energy blending of whole mushroom fruiting bodies exceed those of ChNF sheets but have comparable moduli, which also compares favorably to the strength of CNF nanopapers. In contrast to cellulose nanopapers, chitin nanopapers or sheets prepared from ChNCs are much more hydrophobic probably because of a hydroxyl group in each polymer repeat unit is replaced with an acetyl amine moiety and associated hydrophobic impurities. Advancing water contact angles measured on BC and CNF vary between 11° to 46° , $^{269-272}$ while those measured on chitin nanopapers were 50° to 55° .^{66,192} Static advancing water contact angles on hot compacted ChNC sheets even reached 87° ,²⁷³ and those for FChNF films ranged from 87° to 122° ,^{258,274} because of the presence of nonpolar impurities such as alcohols and acid derivatives, which are also responsible for the aroma of fungi. However, after further purification of fungi extracts, the static advancing contact angles decrease to 80°,258 approaching those of crustacean chitin sheets. The reduced hydrophilicity of various chitin nanopapers and films could be an advantage for instance for packaging applications.

Chitin nanopapers, sheets, and films have been explored for various applications, such membranes and filters and potential packaging applications.²⁷⁵ Unfortunately, not much is known about the barrier properties of pure chitin nanopapers, sheets, or FChNF papers; however, their moduli and strengths are promising, albeit the strains to failure and reported works of fracture (related to the fracture toughness) are still quite low.

Fungal Chitin in Biomedical Use. Fungal material has been used globally for medical applications since ancient times, where it was used as a styptic to stop bleeding and as a crude precursor to modern antibiotics for treatment of infections.^{276–279} However, it was not until the 1970s that true medical materials produced from fungi were investigated. Prudden worked extensively on powdered fungal mycelium as a topical agent for accelerating wound healing and found that both untreated and NaOH or HCl treated mycelium improved the tensile strength of wounded skin, a result that was reproducible using crustacean chitin.^{280,281} Fungal chitin has also been linked to proliferation of fibroblasts and keratinocytes, which are important for creating a new skin base layer, and the activity of matrix metalloproteinases (MMPs) (human cells).^{282,283} In 1997, a research group from Taiwan extracted a chitin-polysaccharide mixture from Ganoderma tsugae, comprising β -1–3-glucan (~60%) and Nacetylglucosamine (\sim 40%), which was used to create a weaveable skin substitute called Sacchachitin. This novel wound dressing was tested on rats²⁸⁴ and on guinea pigs²⁸⁵ before being tested in a preliminary clinical trial on two human patients with chronic wounds in 2005.²⁸³ The animal studies showed that Sacchachitin improved wound healing significantly compared with gauze and had comparable performance to Beschitin, a commercially available wound dressing from crustacean chitin developed in 1988. Improvements in healing were also observed in human trials. However, the surge of interest into the more promising medicinal properties of chitosan in the 1980s and its potential for drug delivery systems resulted in a research swing, resulting in a dearth of further literature utilizing fungal chitin for production of medical materials.²⁸⁶⁻²⁹³ Readers interested in the medicinal properties of chitosan and drug delivery systems are directed to comprehensive reviews on these areas.²⁹⁴⁻²⁹⁹

Mycelium Foams for Packaging Applications. Up to this point, we have described chitin-based materialscomposites and nanopapers-that hold just a promise for authentic applications without any large-scale industrial commitment. By contrast, mycelium foams introduced in this section are genuinely used as packaging materials by real companies. The driver for their use is that packaging materials typically do not have many material property requirements, other than providing mechanical protection to consumer goods and being low in density to minimize shipping costs. Indeed, one of the key challenges in replacing plastic packaging materials is competing with their low cost. Here, mycelium can offer an alternative packaging foam based on its native construction. So far in this Perspective, most of the cited accounts have utilized the fruiting body of mushrooms-the portion above ground (Figure 4b)—as the source for chitin. Mycelium, on the other hand, is the filamentous root-like growth of mushrooms and other fungi, containing mainly chitin and β -glucan, yielding a natural nanocomposite architecture associated with impressive mechanical properties.^{258,300} Fungal mycelium growth can be used to bind lignocellulosic matter in a natural, low energy, self-assembling

Biomacromolecules

production process occurring at ambient temperatures, which also sequesters carbon.^{301,302} The use of mycelium as a binder is not entirely new with some Asian cultures having utilized solid-state fermentation of *Rhizopus oligosporus* to bind soybeans into solid cakes for centuries, producing foods such as Indonesian tempeh. Applying this principle allows for the upcycling of agricultural residues, which have almost no value of their own, into porous, low-cost, highly sustainable, and fully biodegradable composite materials resembling polystyrene foam (Figure 10).³⁰¹ Such materials are already available as commercial products.



Figure 10. Mycelium composite foams used to package consumer products. Reprinted by permission from Ecovative Design LLC, Copyright 2019.

With the main feedstocks for mycelium composite production being agricultural wastes and byproducts, such as corn stalks, cotton carpel, and rice hulls, and the negligible cost associated with the natural manufacturing process, these composites can have raw material costs as low as 13 \$US/m³ for rice-hull-based mycelium composites (Table 6).301,303,304 This makes their raw material costs lower than expanded polystyrene, which uses the styrene monomer as a precursor material, and their biological manufacturing process much less energy intensive.³⁰³ Mycelium composites are also very lightweight with typical densities of approximately 100 kg/ m^3 , but as low as 59 kg/m³, which is only slightly heavier than expanded polystyrene, which typically has densities of up to 50 kg/m 3 . 302,304,305 In fact, mycelium composites exhibit properties making them ideal for packaging applications with elastic moduli, dimensional stability, degradation rates, flame retardance characteristics, and thermal conductivity, all within acceptable limits for packaging applications.³⁰¹ However, the key advantage of mycelium composites occurs at the end of their life, with simple and sustainable disposal possible through normal household garden composting.

Table 6. Cost (US/m^3), Density (ρ , kg/m³), Compressive and Flexural Strength ($\sigma_{compressive/flexural}$, kPa) of Mycelium Composites for Packaging Solutions⁴⁴

material property	unit	expanded polystyrene	mycelium foam ^b
raw material cost	\$US/ m ³	18-83 ^c	13-61
density, ρ	kg/m^3	11-50	59-224
compressive strength, $\sigma_{ m compressive}$	kPa	33-690	1-490
flexural strength, $\sigma_{ m flexural}$	kPa	70-696	7-220

^{*a*}Data from ASTM International,³⁰⁵ Jones et al.,³⁰⁶ Jones et al.,³⁰⁷ Holt et al.,³⁰⁸ Travaglini et al.³⁰⁹ Appels et al.³¹⁰ Commercial values reflect the most recent available literature and are adjusted for inflation to the 2019 \$US value. ^{*b*}Based on compositions of 25 wt % wheat grains and 75 wt rice hulls (59 kg/m³) or 100 wt % wheat grains (224 kg/m³) with values of 37 (rice hulls) and 170 (wheat grains) \$US/tonne. ^{*c*}Based on a cost of \$US 1660 per metric ton of styrene monomer in the United States in September 2019 and a polystyrene density of 11–50 kg/m³.

Although mycelium composites have been considered a viable replacement for packaging materials, such as expanded polystyrene, for over 10 years their adoption has been slow. Dell currently uses mycelium composites for packaging business servers,³¹¹ and IKEA has suggested that they will also adopt the material for packaging their products.³¹² In addition to packaging applications, mycelium composites have also received significant academic and commercial interest for use in construction and architecture, with their low thermal conductivities, significant fire resistance, and outstanding acoustic absorption properties making them excellent insulation and partitioning materials.

■ FUTURE PERSPECTIVES

It is evident from this Perspective that renewable nanomaterials, such as cellulose nanocrystals, nanofibrillated and bacterial cellulose, 5,313 chitin, 314 and starch, 315 have attracted significant research interest and have been explored for numerous applications—a trend that has been growing within the past 10–12 years. In particular, these nanomaterials are promising reinforcements for (renewable) nanocomposites 11,316 because of the possibility of exploiting their stiffness and strength. In light of the recent quest to replace "single-use plastic" with renewable alternatives, chitin is one option 317,318 currently being (re)explored (it was already explored until the 1940s, prior to the development of oil-based synthetic polymers²⁵⁵).

In this Perspective, we have discussed the potential applications of ChNF and ChNC as renewable reinforcements for polymers to produce sustainable chitin nanocomposites. The tensile moduli and strengths of chitin nanocomposites are partially on par with commodity polymers (PET, PP, and HDPE). To date, we have demonstrated that high-performance chitin-reinforced polymer composites only exceed the mechanical performance of common bulk polymers if the chitin loading fraction is greater than 50 wt %. However, tensile modulus and strength of a material are only half the story. Fracture toughness is another important but often neglected parameter concerning biobased composite design. To evaluate the potential of chitin nanocomposites as alternatives to PET, PP, and HDPE, their fracture resistance and damage tolerance should be quantified.

In addition to their role as polymer reinforcement, ChNC and ChNF can also be directly used to produce self-standing sheets or nanopapers. We demonstrate in this Perspective that chitin nanopapers and films could play a role in the quest to replace "single-use plastics" but only if fracture toughness can be further improved. There are many reasons to explore the use of fungi and their cell walls as major potential sources of high-performance nanofibers. These FChNFs are of natural origin, are very easily extracted from fungal biomass (you just need to "get an old blender"),¹⁷⁶ and are effectively "prepregged" by nature, with mild extraction conditions ideal for the manufacture of FChNF films (or nanopapers). These films are true bionananocomposites; all components, that is, reinforcement and matrix, are produced by nature, and the matrix is even grafted to the reinforcement. FChNFs contain varying matrix quantities and polymer types (i.e. glucan or chitosan), depending on the species from which they were extracted, which eases film (or nanopaper) formation. However, unfortunately this means that FChNFs rarely contain pure chitin. Although this might be a disadvantage for some applications, it is advantageous if ChNCs are of interest as fungal chitin nanocrystals (FChNC), which do easily disperse to form stable colloidal suspensions.¹⁷

The tensile properties of FChNF films depend on the composition of the FChNF, which in turn depends on the type of fungal species and region within the fungus. The tensile properties of chitin nanopapers and films are on par with CNF nanopapers, although their elastic moduli are significantly lower. FChNF films offer a boarder property envelope than ChNF nanopapers. Barrier properties of polymer materials, but especially of polymer films, are always important.³¹⁹ Unfortunately, limited literature exists documenting the barrier properties of pure chitin nanopapers²⁶⁰ and nothing is yet known about the barrier properties of FChNF films. It has been shown that chitin nanopapers have oxygen and CO2 barrier properties exceeding those of PET; however, those properties were only determined at 0%RH.²⁶⁰ Furthermore, it has been shown that spray coating polylactic acid (PLA) with alternating layers of CNCs and ChNF results in significant reductions in the oxygen permeability of this composite structure as compared to neat PLA. These reductions are even present at 70%RH, but unfortunately, the water vapor transmission rate is still controlled by the PLA substrate layer.³²⁰ Alternative solutions to address the challenges related to water vapor barrier properties have yet to be identified.

Considering that chitin (both of crustacean and fungal origin) can be used as reinforcement for polymers as well as self-standing sheets or films, the comparative mechanical performance of these materials with respect to each other is of interest. Figure 11 summarizes the tensile properties of chitin nanocomposites, chitin nanopapers, and ChNC sheets, in addition to FChNF films. It can be seen that chitin nanocomposites (as documented in research conducted to date) generally have poorer performance than well-consolidated chitin nanopapers, ChNF films, and even ChNC sheets. Many readers would argue that a major hurdle in the use of ChNF papers, ChNC sheets, or FChNF films for various advanced engineering applications is the sheet thickness, which is currently limited to <0.2 mm. We have made initial attempts to produce thicker samples of FChNF films (which could also be called in analogy to thick polymer films: sheets) with a thickness of about 1 mm and an areal density of 1000 g m^{-2} produced by simple vacuum assisted dewatering of A. bisporus whole mushroom extract with a consistency of 0.8 w/v%, followed by press-forming (Figure 12). The resulting FChNF



Figure 11. Comparison of the tensile properties of ChNF/ChNC reinforced polymer nanocomposites, regenerated chitin films, chitin (ChNF) nanopapers, sheets of chitin nanocrystals (ChNC), and fungal chitin nanofiber (FChNF) films as well as a FChNF sheet (shown in Figure 12).

sheet possessed a modulus and strength of 7.5 GPa and 110 MPa, respectively. This could potentially open up new applications for fungal-based chitin nanomaterials in "paper molding" applications.

Many fungal species can also be grown readily in industrialscale liquid fermenters and the mycelium harvested efficiently (think myco-protein, which is sold as meat substitute; Quorn being a leading example). Moreover, fungi are nature's most effective solid-state decomposer organisms, able to explore and assimilate complex 3D substrates (e.g., wood and agricultural waste, such as straws, etc.), other recalcitrant materials, municipal solid waste (think composting), and even several synthetic polymers (e.g., PLA) under particular circumstances and convert them into this valuable fungal cell wall material. This cell wall material can be extracted or used as mycelium composite for packaging or architectural applications. Fungi do this without the need for extraneous energy sources (pH and some nutrients must be suitable) commonly under mesophilic conditions. Additional research in mycelium composite material technology is required to address key limitations of these materials including their slow production rate compared with polymer processing and some parameters that fall outside of desirable ranges, such as water absorption. These developments, in addition to the exceptional potential of mycelium composites as a fully biodegradable, cheap, and lightweight alternative to synthetic foams, such as expanded polystyrene, which also upcycle agricultural residue and sequester carbon, suggest that a more rapid adoption of this material platform would be appropriate if environmental sustainability is to be preserved for our children and humankind to come.

These points highlight the potential for highly efficient and environmentally sustainable exploitation of FChNFs for a new generation of materials for a highly efficient and environmentally compatible exploitation of FChNFs in a new generation of materials. The great diversity of fungi (potentially some 5 million species¹⁴⁵ ranging from unicellular to multicellular, mycelium to mushrooms) and their extraordinary range of environmental tolerances (deserts to the deep sea, high salinity, contaminated environments, below 0 °C to thermophiles) opens a huge range of possible



Figure 12. Photographs of (a) a wet filter cake of *A. bisporus* whole mushroom extract with an intended grammage of 1000 g/m² obtained by filtration of a FChNF suspension with a consistency of 0.8 w/v%, (b) wet pressed cake and (c and d) hot pressed cake with a final sheet thickness of 0.7 mm.³²¹

cultivation and processing conditions for systematic, large-scale production.

A method to utilize the potential of chitin is to spin them into macrofibers. Chitin could be dissolved directly in a solvent, such as 5 wt % LiCl/dimethylacetamide or 5 wt % LiCl/*N*-methyl-2-pyrrolidone and spun into fibers. The viscose process used to dissolve cellulose could also be used to dissolve chitin and wet spun into fibers. In fact, these ideas have been explored in the 1970s to produce chitin fibers.³²² Chitin fibers with tensile strengths as high as 350 MPa have been produced. These fibers (and even fungal mycelium)³²³ can then be processed into textile fabrics for the fashion industry (e.g. "trash fashion"), reducing the environmental burden associated with cellulose textile fibers.

Specifically, for fungal chitin, the chitin microfibrils in fungal cell walls are orientated. As of yet, no one has attempted to make use of this inherent orientation of FChNF, which offers an unexplored potential for materials development if fungal cell walls can be processed to preserve this orientation or, alternatively, processed in such a way as to reinstate such orientation possibly via self-assembly mechanisms, which occur in living fungi. It is believed that in forming the cell walls of fungi, a certain amount of templating is established at the cell membrane, with ongoing self-assembly occurring as the microfibrils are synthesized in the cell wall outside the cell membrane (but with the materials production and export from the cell controlled intracellularly). There are then possible/ likely periods of further templating by the cell membrane in order to create "deliberate" reorientation, in addition to cell extension growth, which causes a degree of physical reorientation.

It is our contention that the fungal biorefinery as a source for high-performance material has been overlooked for far too long and that FChNFs will become one major source of advanced reinforcement for sustainable biobased composites in the 21st Century.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.bio-mac.9b01141.

Comprehensive tables listing nanocomposite and nanopaper applications of nanosized chitin (PDF)

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Notes

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ABBREVIATIONS

ChNC, chitin nanocrystal; ChNF, chitin nanofiber; CNC, cellulose nanocrystal; CNF, cellulose nanofiber; DMAc, dimethylacetamide; TEMPO, 2,2,6,6-tetramethylpiperidine-1-oxyl radical

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Biomacromolecules

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